

Pesticide FactSheet

Name of Chemical: Etoxazole

Reason for Issuance: Conditional Registration

Date Issued: August 22, 2002

1. <u>DESCRIPTION OF CHEMICAL</u>

Generic Name: 2-(2,6-difluorophenyl)-4-[1,1-dimethylethyl)

-2-ethoxyphenyll-4,5-dhydrooxazole

CommonName: Etoxazole

Trade Name: TetraSan™ 5 W DG

EPA PC Code: 107091

Chem calAbstracts

Service (CAS) Number: 153233-91-1

YearofInital

Registration: 2002

Pesticide Type: Miticide Ovicide

Chem icalClass: DiphenylOxazoline

U.S. Producer: ValentUSA Corporation

2. <u>USE PATTERNS AND FORMULATIONS</u>

Application Sites: O mam entalplants grown in greenhouses,

shadehouses, and htthhouses.

Types of Formulations: 95.4% technical product

5.0% waterdispersible granular end-use product

Types and M ethods C om pressed air sprayers, hydraulic sprayers

of Application or ground boom sprayers.

Application Rates: An application rate of 8 to 16 ounces of product per

100 galbns ofwater (which equates to 0.025 to 0.05 pounds ofactive ingredient/100 galbns or 16 to 32 ounces/acre). A second application can be applied 14 days after the first application if necessary. No more than two (2) applications per cropping season or two (2) applications in six (6) months are allowed.

Carrier: Water

3. <u>SCENCE FINDINGS</u>

Etoxazole is a memberofthe diphenyloxazoline class of insecticides. Available productchem istry, toxicology, ecological effects and environmental fate data supporting the ornamental use pattern has been reviewed. The data and estimated risks to hum an health and the environment from its use on ornamental plants grown in greenhouses, shade, and lathhouses are summarized below.

ChemicalCharacteristics

Table 1. Chem ical Characteristics

Property	Techn <i>i</i> cal	End-Use
Cobr	Munsellcobrnotation N 9.5 (FG A I) White N 9.5 /with 90% reflectance (PA I)	Brown
PhysicalS tate	Lum py powder (TGAI) Free Flow ing crystalline Powder (PAI)	Brown granules
0 dor	Musty odor (TGAI) No obvious odor(PAI)	N ⁄A
Oxidation/meduction:Chemical incompatiblity	TS was found nothaving any oxidizing properties	No reaction was observed
Flam m ability/Flam e Extension	Non flam m able	N /A
Explodability	Non-expbsive	Notexpbsive
S torage S tability	Stable forone yearunder warehouse conditions	In progress. Study in it is ted 3/9/00.

Property	<u>Technical</u>	End-Use
M iscibility	N 🛦	N 🔼
Comosion Characteristics	In progress. Study in triated 3/9/00.	In progress. Study in itiated 3/9/00.
рН	62	5.94
Melting Point	101.5 to 102.5C (PAI)	N ⁄A
D ensity	37.6 bs/cu.ft.(TGAI) 12389 (Relative density of PAI)	3 4.2 b./⊞ at21 ["] C
Solubility (W ater) Column elution m ethod; shake flask m ethod	3.99 x 10 ⁻⁵ g/lin distilled water at 10 °C 7.04 x 10 ⁻⁵ g/lin distilled water at 20 °C 6.69 x 10 ⁻⁵ g/lin distilled water at 30 °C	N 🕰
VaporPressure	7.0 x 10 ⁻⁶ pascals @ 25 ["] C (PA 1)	N ⁄A

¹ TechnicalG rade Active Ingredient

Toxicology Characteristics

Table 2. A cute Toxicity Data on Etoxazole Technical

Guide ine No.	Study Type	R esu l ts	Toxicity Categor v
OPPTS 870.1100	Acute O ral-M ouse	LD ₅₀ > 5000 m g <i>]</i> kg	N
OPPTS 870.1100	Acute Oral-Rat	LD ₅₀ > 5000 m g/kg	N
O PPTS 870 1200	Acute Dem al-Rat	LD ₅₀ > 2000 m g <i>]</i> kg	ш
OPPTS 870.1300	Acute Inhalation -Rat	LC ₅₀ > 1.09 m g/L	ш
OPPTS 870.2400	Prim ary Eye Imitation -Rabbit	Notan eye <i>irrit</i> ant	IV
O PPTS 870.2500	Primary Skin Imitation - Rabbit	Nota dem alimiant	IV

² Pure Active Ingredient

OPPTS	DemmalSensitization -Guinea Pig	Nota dem alsens itiz er	N /A
870.2600			

Table 3. A cute Toxicity Data on TetraSan™ 5 W DG Insecticide

Table 3. Active TOXILLY Data Off Tedasait 5 W DG IISeculture			
Guide ine No.	Study Type	Results	Toxicity Categor v
OPPTS 870.1100	Acute Oral-Rat	LD ₅₀ = 4274 m g/kg	ш
OPPTS 870 1200	Acute Dem al-Rat	LD ₅₀ > 5000 m g <i> </i> kg	Ŋ
OPPTS 870.1300	Acute Inhalation -Rat	LC ₅₀ > 2.05 m g/L	IV
OPPTS 870 2400	Prim ary Eye Imitation -Rabbit	Conjunctivitis in 3/3 eyes atone hourafter instillation. All imitation resolved by 48 hours.	ш
OPPTS 870.2500	Primary Skin Imitation - Rabbit	Nota dem alimiant	V
OPPTS 870,2600	Dem alSensitization -Guinea Pig	Nota sens it zer	N /A

Table 4. Subchronic Chronic Toxicity Data on Technical Etoxazole

Guide ine No./Study Type	Results (mg/kg/đay)
870.3100 90-Day Feeding Rat	NOAEL = notdeterm ined LOAEL = 300.4/336.6 (M/F), based on clinical signs, clinical chem istry, increased liverweights, and histopathology
870.3100 90-Day Feeding Rat	NOAEL = 183/205 M/F) LOAEL = 183.7/204.8 M/F) based on increase in hepatic enzyme levels, increased liverweights and centribbularhepatocellularswelling in both sexes and liverenlargement in females only
870 3150 90-Day Feeding M ouse	NOAEL = 213.6/250.5 M/F) LOAEL = 878.4/994.5 M/F), based on perportal hepatocellular necrosis, increased alkaline phosphatase levels, accompanied by increased relative liver weight, liveren largement, and centrib bular hepatocellulars welling

Guide ine No /Study Type	Results (mg/kg/day)	
870 3150 90-Day Feeding Dog	NOAEL = 5.33/5.42 (M /F)	
	LOAEL = 53.7/55.9 M/F), based on clinical signs (vom iting foam y fluid and mucous stool), clinical chemistry, increased liverweights, and on centribbular hepatocellularswelling in the liverand acinarcellatrophy in the prostate.	
870 3200 21-Day DermalTox Rat	NOAEL = 1000	
	LOAEL > 1000, no effects noted.	
870.3700 Developm entalToxRabbit	NOAEL = <u>Matemal</u> : 200 <u>Developmental</u> : 200	
	LOAEL = Matemal: 1000, based on liverenlargem entand decreased body weightgains and food consumption Developmental: 1000, based on increased incidences of 27 presacral vertebrae and 27 presacral vertebrae with 13 rbs (skeletalvariations) in the fetuses	
8703700 DevelopmentalToxRat	NOAEL = <u>M atemal</u> : 1000 <u>D evelopm ental</u> : 1000	
	LOAEL = <u>Matemal</u> : notdeterm ined <u>Developmental</u> : notdeterm ined	
870.3800 2-G en R eproduction R at	NOAEL = <u>ParentalSystem ic</u> : 20 <u>Offspring System ic</u> : 20 <u>Reproductive</u> : 100	
	LOAEL = <u>ParentalSystem in the Pand</u> F	
	m ales and increased adrenalweights in the P fem ales <u>Offspring Systemic</u> : 100, based on pup mortality <u>Reproductive</u> : Not determined	
870.4100 1-YearFeeding Dog	NOAEL = 4.62/4.79 M/F) LOAEL = 23.5/23.8 M/F), based on increased alkaline phosphates activity, increased liverweights, liverenlargement (fem ales), and incidences of centribbularhepatocellularswelling in the liver	
870.4200 78 -W eek Carrinogenir M ouse	NOAEL = 241/243 (M /F) LOAEL = Notdeterm ined, No evidence of carchogenicity	
870.4300 2-YearFeed, Carcinogenic Rat, 52-week interin report	NOAEL = 2.05/2.4 M/F) LOAEL = 208/247 M/F) based on increased absolute and relative liverweight and elongation, whitening and abrasion of the incisors in both males and females, Y-glutam yltranspeptidase in males only, and urinary volume, chronic nephropathy, total protein and protein uria in females only	

Carcinogenicity

Classification of the carcinogenic potential of etoxazole is not required at this time as all hum an exposures will be short-term (no more than 2 applications per cropping season or 6-m onth period). Although they are not required at this time, carcinogenicity studies have been submitted for Agency consideration in connection with pending food uses for etoxazole; the pending food uses will be evaluated at a later date. Preliminary results suggest that there is no evidence of carcinogenicity in the ratorm ouse as a result of or administration of etoxazole. The ratand mouse carcinogenicity studies listed in Table 2 are separate studies submitted previously. Although these studies showed no evidence of carcinogenicity up through the highest dose tested, adequately high doses were not tested.

Mutagenicity

The Agency has concluded that there is not a concern form utagenizity resulting from exposure to etoxazole. Six acceptable genetic toxicology studies were available for review. The results from these studies indicate that etoxazole was not mutagenic in Salm one la typhim urium, Escherichia coli, and L5178Y TK +/- mouse lymphom a cell assays. There was also no evidence of clastogenizity in vitro, and etoxazole gave a negative response for the induction of micronucleated polychromatic erythrocytes in bone manow. Also, etoxazole did not induce unscheduled DNA synthesis. O verall, the data suggest that etoxazole is negative form utagenizity in vitro and in vivo.

Toxicological Endpoints

The dose atwhich no adverse effects are observed (the NOAEL) from the toxicobgy study identified as appropriate foruse in risk assessment is used to estimate the toxicobgical levelofconcern (LOC). The bwestdose atwhich adverse effects of concern are identified (the LOAEL) is sometimes used for risk assessment if no NOAEL was achieved in the toxicobgy study selected. An uncertainty factor (UF) is applied to reflect uncertainties inherent in the extrapolation from laboratory animal data to humans and in the variations in sensitivity among members of the human population as well as other unknowns. An UF of 100 is routinely used, 10X to account for interspecies differences and 10X for intraspecies differences.

The acute, sub-chronic and chronic (non-cancer) toxicological endpoints that have been established for etoxazole are sum marized in the Table 5.

Table 5. Acute, Sub-chronic and Chronic (non-cancer) Toxicological Endpoints

Exposure Scenario	Dose (mg/kg/day) UF/MOE	Hazard Based SpecialFQPA Safety Factor	EndpointforR isk Assessm ent
	Die	etary R isk Assessm e	ents
A cute D ietary	N /A Non-food use on l	7.	
Chronic Dietary	N/A Non-food use only	7.	
IncidentalOral	N/A Noresidentialus	es.	
	Non-Dietary Risk Assessments		
Derm al Alldurations (1 day \$ 6 m onths)	No hazard quantitation required.		28-Day Derm alStudy-Rat NOAEL = 1000 m g/kg/day No system is effects noted up to 1000 m g/kg/day.
Residential	n ⁄A	n A	
0 ccupational	n 🛦	N/A	
Inhalation Alldurations ³ (1 day \$ 6 m onths)	Oral NOAEL = 4.62		Chronic OralToxicity Study -Dog LOAEL=23.5 mg/kg/day based on increased alkaline phosphatase activity, increased liverweights, liverenlargement
Residential	и ∕А	N /A	(fem ales), and incidences of centribbular hepataocellularswelling in the liver
0 ccupational	100	N A	
Cancer	N /A Non-food use only.		

³ Since an orallendpointwas selected, an inhalation factor of 100% should be used in route-to-route extrapolation.

Hum an Exposures and Risks

Acute and Chronic Dietary Risk: Notapplicable, since there are no food uses of etoxazole.

Occupational Risk: Estimated margins of exposure $(M \circ E)$ for all handlers greatly exceed the target $M \circ E$ of 100 in all cases provided handlers wearbaseline personal protective equipment (PPE) of brighted shirt, brighted handlers wearbaseline personal protective equipment (PPE) of brighted shirt, brighted handlers socks and a 12-hour restricted entry interval (REI) is observed. The end-use label includes both requirements. Two handler occupational activities are indicated from the use of etoxazole, mixing and bading the pesticide and applying the pesticide. The end-use product is packaged in water-soluble bags thus greatly reducing exposure during mixing and bading (estimated $M \circ E = 2,700,000$). Although many large 'houses" utilize automated spray systems whereby no individual is involved in the actual spraying procedure, and the end-use label allows for no less than a 25-gallon tank mixture, $M \circ E$ swere calculated for applicators

using bw-and high-pressure hand-held spray wands as the application scenarios of highest exposure. The MOE for the bw-pressure hand-held spray wand is estimated to be 1,700, the MOE for the high-pressure hand-held spray wand is estimated to be 8,000. Thus, the Agency's evelof concern is not exceeded for any application method. Post-application exposure to re-entry workers was not assessed because no dermaltoxicity endpoints were identified. A restricted entry interval of 12 hours is required.

Environm entalCharacteristics

Available environmental fate data indicate that etoxazole degrades atm oderate rates at pH 5 (approximately 10 days), and it is relatively stable between pH 7 and 9. Etoxazole showed moderately rapid biodegradation in a variety of soils ranging from a variety of characteristics, a total of seven values available yielded a mean half-life of 20 5 days. Etoxazole is immobile in several soils tested, with $K_{\rm foc} > 5000$ in seven out of eight soils tested. There appears to be little potential for etoxazole to be transported with water, although transport of residues adsorbed to the eroding soil is possible. Based on etoxazole's greenhouse, shadehouse, and lathhouse use sites, environmental exposure should be limited.

EcologicalCharacteristics/Risk

Available ecological effects data indicate that etoxazole is practically non-toxic to birds in both acute oralbasis ($LD_{50} > 2000 \, \text{mg/kg}$) and sub-acute basis ($LC_{50} > 5000 \, \text{ppm}$). Based on etoxazole's use sites, significant exposure to non-target organisms is not expected to occur. Therefore, chronic testing was not required.

In contrast, etoxazole is considered very highly toxic to aquatic invertebrates in acute testing. Acute toxicity tests of etoxazole with fieshwater fish are considered invalid for reasons including solubility problems and failure to use flow-through testmethods, among others. Additional studies are required to characterize acute toxicity on fieshwater fish. As with avian risk, however, significant exposure to aquatic, non-target organisms is not expected to occur. Therefore, chronic testing is not required.

4. <u>SUMMARY OF REGULATORY POSITION AND RATIONALE</u>

Available data provide adequate information to support the conditional registration of etoxazole technical and end-use products for use on ornam entalplants grown in greenhouses, shadehouses, and lathhouses.

Use, Form ulation, Manufacturing Process or Geographic Restrictions

<u>Use Directions - General Precautions</u>

Do not apply through any type of irrigation or chem igation system.

Do notapply this product in a way that will contact workers or other persons, either directly or through drift. Only protected handlers may be in the area during application.

Do notenterorallow workerentry into treated areas during the restricted entry interval (REI) of 12 hours.

Do not contam inate water, food or feed by storage or disposal.

Do not contam in ate waterwhen disposing of equipm entwashwaters.

Use Directions - Ornam entals Grown in Greenhouses, Shadehouses, and Lathhouses

Apply no more than two times percropping cycle.

Apply no more than two times persix months.

5. SUMMARY OF DATA GAPS

Comosion characteristics (Guideline 830.6320) and storage stability (Guideline 830.6317) on technical etoxazole and end-use product

Freshwater fish acute toxicity (Guideline 850 1075) on technical etoxazole

90-Day subchronic inhalation study on rats (Guideline 870 3465) on technical etoxazole

6. CONTACT PERSON AT EPA

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